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Triple Junction Segregation Dominates the Stability of Nanocrystalline Alloys

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and TJs. Our work highlights the need to account for TJ segregation effects in order to understand and predict the evolution of nanocrystalline alloys under extreme environments.

KEYWORDS: *triple junction, solute segregation, nanocrystalline materials, atomistic simulations, grain boundary*

Nanocrystalline (NC) materials are characterized by their
nanometer-sized crystalline grains that are internally
initial through a natural of interfaces termed grain joined through a network of interfaces, termed grain boundaries (GBs), and line defects, termed triple junctions (TJs), where three GBs meet. Owing to their small grain size, NC materials exhibit unique combinations of functionalities including mechanical strength,^{[1](#page-7-0)} wear resistance, $2^{2,3}$ transport properties,^{[4](#page-7-0),[5](#page-7-0)} and electrochemical performance.^{[6](#page-7-0),[7](#page-7-0)} However, the high density of GBs in NC materials renders them thermally unstable with respect to grain coarsening and concomitant evolution of their crystal-size dependent proper-ties even at low homologous temperatures.^{[8](#page-7-0),[9](#page-7-0)} Mitigating grain growth during synthesis and processing treatments or under operating environments remains one of the main hurdles to the use of NC materials in a wide range of engineering applications.^{[10](#page-7-0)}

Much of the research dealing with improving the thermal stability of NC metals has been geared toward employing GB solute segregation to mitigate boundary motion.^{[11](#page-7-0),[12](#page-7-0)} With this approach, the NC material is alloyed with an element that preferentially segregates to GBs, which then slows boundary migration and grain growth through various mechanisms. $11-15$ $11-15$

While several studies demonstrated thermally stable NC materials through alloying and subsequent GB segregation,[15](#page-7-0)[−][22](#page-7-0) absent from such studies is a detailed understanding of the role of TJ solute segregation in the evolution of nanostructures and stability of NC alloys. In addition to GBs,

dislocations, and point-defects, TJs are an important part of the repertoire of defects in crystalline solids. Indeed, numerous studies have demonstrated that TJs influence a host of materials phenomena and properties, including diffusion, $23,24$ thermoelectric^{[25](#page-7-0)} and mechanical^{26−[28](#page-7-0)} properties, and GB dynamics.[29](#page-7-0)−[33](#page-7-0) To demonstrate the increased contribution of TJs with decreasing grain size, one considers a two-dimensional representation of a NC material with a characteristic grain size *L* (see [Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf) Figure S-1). For this geometry, the TJ density ρ_{T1} , in units of number of TJs per unit area, and GB density $ρ_{GB}$, in units of length of GB segments per unit area, are related through (see the [Supporting](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf) [Information](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf))

$$
\frac{\rho_{\rm TJ}}{\rho_{\rm GB}} \propto \frac{1}{L} \tag{1}
$$

[Figure](#page-2-0) 1a is a schematic plot of eq 1 depicting values of $\rho_{\text{TI}}/$ ρ _{GB} for micrometer- and nanometer-size crystalline grains, where a drastic increase in TJ density relative to the GB

Figure 1. (a) Schematic plot of [eq](#page-1-0) 1 showing values of ρ_T/ρ_{GB} for micrometer- and nanometer-size crystalline grains. For an annealed Pt-10 at. % Au alloy thin film: (b) a representative STEM-HAADF image depicting nanocrystalline grains, (c) quantified Au atomic concentration map of the structure in (b), and (d) an enlargement of the region marked by the dashed line in (c), highlighting increased Au segregation at several TJs. See the Supporting [Information](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf) for details on sample preparation and data acquisition.

Figure 2. (a) Grain structure of a pure NC Pt with a [111] crystallographic direction for the out of plane direction. For a Pt-10 at. % Au alloy, close-up views showing representative (b) unrestricted segregation in [001] and (c) TJ segregation in [111] NC systems. (d) For the [111] NC Pt-10 at. % Au alloy, a plot of the rescaled total energy *E*̅(*t*) during MC-MD, where chemical equilibrium is inferred from the asymptotic behavior of *E*̅ curves. In (a−c), nanograins are assigned unique colors, GB atoms are in black, and Au atoms are in yellow.

density is seen as the grain size *L* is reduced into the nanoscale. Furthermore, since TJs exhibit atomic structures that are distinct from the GBs they are connected to, it is natural for solute atoms to have interactions with TJs that are different than those with GBs.^{[34](#page-7-0)−[37](#page-7-0)} Indeed, recent experimental studies revealed solute enrichment at TJs in NC alloys.[22](#page-7-0),[35,38](#page-7-0)−[40](#page-8-0) As a demonstration, Figure 1 shows results for a Pt-10 at. % Au NC alloy, where Figure 1b depicts an image using transmission electron microscopy (TEM) of the nanograin structures in this NC alloy. Figure 1c shows the concentration map, obtained using energy-dispersive X-ray spectroscopy (EDS), of the structure shown in Figure 1b, and Figure 1d is a close-up view of the region marked by the dashed line in Figure 1c, clearly showing strong Au enrichment at TJs in addition to GB segregation. Owing to the geometric degrees of freedom describing GBs and $T\int_{s}^{41,42}$ $T\int_{s}^{41,42}$ $T\int_{s}^{41,42}$ variations in solute segregation

levels at these defects can be seen in Figure 1c. The NC Pt−Au alloy thin film was prepared, annealed, and charcterized as discussed in ref [43](#page-8-0) (see the Supporting [Information\)](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf). The experimental observations depicted in Figure 1b−d for the NC Pt−Au alloy highlight the need to delineate the role of TJ solute segregation in GB dynamics and grain coarsening in NC alloys.

In this work, we report results of large-scale atomistic modeling studies employing molecular dynamics and Monte Carlo simulations to examine the impact of TJ segregation on the thermal stability of NC alloys. The NC Pt−Au alloy is used as a model system, as Au enrichment to Pt TJs has been experimentally observed here (see Figure 1b−d) and elsewhere. $22,38$ Our analysis clearly demonstrates that TJ segregation plays a dominant role in controlling GB dynamics by locking the GB network and limiting its evolution.

Figure 3. Evolution of the grain structures after 20 ns of annealing at 900 K for the (a) pure NC Pt system and (b, c) [111] Pt-10 at. % Au alloys with (b) unrestricted and (c) TJ segregation cases. (d) For the systems depicted in $(a-c)$, temporal evolution of the average grain size. In $(a-c)$, Au atoms are labeled in yellow, nanograins are assigned unique colors, GB atoms are in black, and the scale bars represent 20 nm.

The atomistic simulations are performed using the LAMMPS software package, 44 and visualization of the atomic structures is done using Ovito.^{[45](#page-8-0)} We use the interatomic potential by O'Brien et al., 46 which is fit to capture many of the thermo-physical properties of the Pt−Au alloy across the Au composition space. First, polycrystalline atomistic systems with a wide range of GB types and Au concentrations are generated using the approach outlined in ref [47](#page-8-0), which employs grain structures generated from mesoscale models to construct periodic atomistic Pt thin films with columnar grains. Here, the *x* and *y* directions define the in-plane coordinates, and the *z* direction describes the out-of-plane axis. We explore cases with [111] and [001] crystallographic directions for the NC film out-of-plane *z* axis, where crystalline grains have random inplane orientations about this common *z* axis. The overall size of the NC systems is approximately $110 \times 110 \times 2.4 \text{ nm}^3$, resulting in a total number of atoms of approximately 1.25 million atoms. Also, for all systems explored in this work the initial average grain size and total number of grains are ∼5 nm and ∼300, respectively. [Figure](#page-2-0) 2a shows a representative NC Pt system with [111] crystallographic direction for the out-ofplane axis, where the nanograins are assigned unique colors and atoms with non-FCC ordering (i.e., GBs) are labeled in black. In order to examine Au concentration effects, we randomly assign Au atoms to yield NC Pt−Au alloys, where we explore cases with nominal Au concentrations of 1, 5, and 10 at. %.

After the initial construction, the hybrid Monte Carlo (MC)−molecular dynamics (MD) scheme^{[48,49](#page-8-0)} is used to equilibrate the atomic structures and compositional degrees of freedom of the NC systems. An MC step is performed with conserved atomic swaps, in which two atoms are randomly selected, and if they are different chemical species, their chemical identities are swapped and accepted with a Boltzmann factor. This MC simulation samples the equilibrium canonical ensemble in a closed system with a fixed number of Pt and Au atoms. In this work, each MC-MD cycle included 500 MC steps followed by a 10 ps MD simulation at 77 K and zero pressure, leading to a trajectory in-phase space that is consistent with the NPT ensemble. This cycle is repeated until the total MC-MD simulation time is 15 ns. In all our MC-MD simulations, a temperature of 77 K is used, as it models cryogenic materials processing routes (e.g., ball milling) commonly used to fabricate NC metallic alloys.^{[50,51](#page-8-0)} Once generated and chemically equilibrated using MC-MD, the NC Pt−Au systems are thermally annealed for 20 ns at zero pressure and 900 K, a temperature corresponding to approximately 40% of the solidus temperature for the Pt-10 at. $%$ Au alloy.^{[52](#page-8-0)} This high-temperature anneal promotes curvature-driven GB migration where the driving force is related to the boundary's local curvature. 53

To reveal and quantify the impact of TJ segregation on GB dynamics and the evolution of nanostructures, we consider two solute segregation scenarios for Au. In the first, we allow the

Figure 4. Demonstration of how TJ segregation locks the GB network. Snapshots depicting the atomic structure and local GB network near several TJs in (a) [001] Pt-5 at. % Au and (b) [111] Pt-10 at. % Au NC alloys. For a better visualization of the dynamics, the bulk atoms are not displayed and Au atoms are labeled in yellow. The dynamics of one GB are revealed by overlaying boundary atoms at different simulation times and assigning them unique colors. Atoms colored in gray correspond to other GBs connected to the TJs.

MC-MD cycles to take place to any pair of Pt−Au atoms regardless of their position within the NC system. We refer to these cases as *unrestricted* segregation, which mimic solute segregation behavior in experimental NC alloys such as the one depicted in [Figure](#page-2-0) 1b−d. In the second scenario, we only allow Au segregation to TJ regions. This constraint is achieved by defining cylinders of diameter ∼12 Å centered around the TJ lines and restricting the MC-MD swaps to only these TJ regions. We refer to these cases as *TJ segregation*. As a temperature of 77 K is used during the MC-MD stage, the evolution of the TJs is limited during the chemical equilibration step. Again, all our MC-MD studies, including TJ segregation cases, are performed at 77 K and zero pressure. For a Pt-10 at. % Au NC alloy, [Figure](#page-2-0) 2b,c shows respectively close-up views of representative cases of unrestricted Au segregation in a [001] NC alloy and TJ segregation in a [111] NC system, where Au atoms are labeled in yellow, nanograins are assigned unique colors, and GB atoms are depicted in black. To ensure that our MC-MD simulations lead to NC alloys approaching chemical equilibrium, we monitor the rescaled total energy \overline{E} of a NC system given by

$$
\overline{E}(t) = \frac{E(t) - E(t = 0)}{E(t = 0) - E(t = 15 \text{ ns})}
$$
\n(2)

where a plateau of \overline{E} vs time (i.e., energy is not changing with time) indicates equilibration of the NC alloy chemistry. For example, [Figure](#page-2-0) 2d shows plots of $\overline{E}(t)$ for the [111] NC Pt-10 at. % Au alloy using both unrestricted and TJ segregation cases, where the asymptotic behavior of $E(t)$ can be seen indicating chemical equilibrium.

The aforementioned approach of generating and annealing NC Pt−Au systems with [111] and [001] textures and three Au concentrations (i.e., 1, 5, and 10 at. % Au) using unrestricted and TJ segregation cases is intended to be representative of the temperature, Au compositions, and texture observed in recent experimental studies on the thermal stability of NC Pt−Au alloys.^{[22,38](#page-7-0)} While we acknowledge that the atomistic systems with unrestricted segregation are more representative of experimentally observed segregation behavior in NC alloys, it is important to emphasize that our computational studies of systems with TJ segregation provide the means to quantitatively describe the relative role of doped TJs in the thermal stability of NC alloys. Also, our computational approach generates crystalline grains with random orientations about the common out-of-plane axis, resulting in GBs and TJs with geometries different from the ones in the experimental NC Pt−Au system in [Figure](#page-2-0) 1. These variations in GB and TJ geometries result in differences in solute segregation levels between our atomistic simulations and the experimental data in [Figure](#page-2-0) 1. Furthermore, while the average grain size in our atomistic simulations is smaller than one in the experimental NC Pt−Au shown in [Figure](#page-2-0) 1b−d, it is important to note that NC materials with an average grain size of less than 10 nm have been synthesized in a wide range of alloys.[54](#page-8-0)−[56](#page-8-0)

We begin with comparing the evolution of grain structures in NC systems, specifically pure Pt and Pt−Au alloys with unrestricted and TJ segregation cases. [Figure](#page-3-0) 3a−c shows snapshots of the [111] NC structures after 20 ns of thermal annealing at 900 K, where pure Pt [\(Figure](#page-3-0) 3a), Pt-10 at. % Au with unrestricted segregation [\(Figure](#page-3-0) 3b), and Pt-10 at. % Au

Figure 5. For [111] NC Pt−Au films with various Au concentrations, a comparison of the role of TJ segregation in grain coarsening relative to that of unrestricted segregation. Snapshots of the grain structures with (a1−a3) unrestricted and (b1−b3) TJ segregation after annealing at 900 K for 20 ns. Alloys with an Au concentrations of (a1, and b1) 1 at. %, (a2, b2) 5 at. %, and (a3, b3) 10 at. % are shown. Temporal evolution of the (c) average grain size and (d) total number of grains for pure Pt and Pt−Au alloys with 1 and 10 at. % Au. In (a) and (b), Au atoms are labeled in yellow, nanograins are assigned unique colors, GB atoms are depicted in black, and the scale bars represent 20 nm.

with TJ segregation only [\(Figure](#page-3-0) 3c) are shown. The initial grain structure for these systems is depicted in [Figure](#page-2-0) 2a. In [Figure](#page-3-0) 3a−c, Au atoms are labeled in yellow, nanograins are assigned unique colors, and GB atoms are depicted in black. NC alloys with unrestricted ([Figure](#page-3-0) 3b) and TJ segregation ([Figure](#page-3-0) 3c) show enhanced thermal stability compared with the pure Pt system ([Figure](#page-3-0) 3a), which experiences rapid grain coarsening. This is shown quantitatively in [Figure](#page-3-0) 3d, which depicts plots of the average grain size as a function of annealing time. The pure Pt system undergoes rapid grain coarsening, where the grains more than doubled in size over 20 ns of annealing at 900 K. This result is in contrast to the cases with TJ and unrestricted segregation, which show a near-flat profile for the average grain size as a function of time. A close examination of [Figure](#page-3-0) 3b,c reveals that the NC alloy with TJ segregation exhibits grain structures that are similar in size to the one with unrestricted segregation. Indeed, [Figure](#page-3-0) 3d shows that after 20 ns of thermal annealing, the average grain size in the NC system with TJ segregation is ∼6.5 nm compared to ∼6 nm in the system with unrestricted segregation and ∼10 nm in pure Pt. The initial average grain size for all NC systems depicted in [Figure](#page-3-0) 3 is ∼5 nm. The results depicted in [Figure](#page-3-0) 3 isolate contributions to thermal stability due to TJ segregation, and it quantitatively demonstrates that doped TJs play an effective role in mitigating grain coarsening in NC alloys.

To reveal how TJ segregation hinders GB migration, we now direct our attention to [Figure](#page-4-0) 4, which depicts representative atomic structures and local GB networks near several TJs for [001] Pt-5 at. % Au [\(Figure](#page-4-0) 4a) and [111] Pt-10 at. % Au ([Figure](#page-4-0) 4b) NC alloys. For a better visualization of the structures, the bulk atoms are not displayed and Au atoms in the TJs are labeled in yellow. Furthermore, [Figure](#page-4-0) 4 depicts the dynamics of one GB by overlaying boundary atoms at different simulation times and assigning each unique colors. Atoms colored in gray correspond to other GBs connected to the depicted TJs in [Figure](#page-4-0) 4. It can be clearly seen that Au segregation to TJs renders these line defects immobile, and as a result, the GBs connected to these TJs evolve by flattening their profiles over time, as depicted in [Figure](#page-4-0) 4. Any further migration of these GBs requires the evolution of the TJs to which they are connected, which is hindered due to TJ segregation. This TJ segregation-induced locking of the GB network and flattening of GB profiles shown in [Figure](#page-4-0) 4 has been observed in all atomistic systems employed in this work (see [Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf) Figure S-2 for more examples). While many studies attributed the experimentally observed thermal stability of NC alloys to GB solute segregation and associated mechanisms of energy reduction of the boundary, and thus driving force for grain coarsening^{11-[13](#page-7-0),[15](#page-7-0)} and GB solute $\text{drag}^{14,57,58}$ $\text{drag}^{14,57,58}$ $\text{drag}^{14,57,58}$ $\text{drag}^{14,57,58}$ $\text{drag}^{14,57,58}$ the results depicted in [Figures](#page-3-0) 3 and [4](#page-4-0) clearly demonstrate that TJ segregation mitigates grain coarsening in NC alloys by locking the GB network, thereby hindering the migration of GBs.

Finally, we investigate the impact of Au concentration levels on the relative roles of unrestricted and TJ segregation in grain coarsening of NC Pt−Au systems. We employ alloys with Au concentrations of 1, 5, and 10 at. % and consider NC systems with $[001]$ and $[111]$ grain orientations. Figure 5 depicts final structures after 20 ns of thermal annealing at 900 K for [111] NC Pt−Au films with various Au concentrations using both unrestricted and TJ segregation scenarios. Panels in the top row (Figure 5a1−a3) depict unrestricted segregation cases, while panels in the bottom row (Figure 5b1−b3) show systems with TJ segregation only. Pt−Au alloys with an Au concentration of 1 at. % (Figure 5a1,b1), 5 at. % (Figure 5a2,b2), and 10 at. % (Figure 5a3,b3) are shown. In all these

panels, Au atoms are labeled in yellow, nanograins are assigned unique colors, and GB atoms are depicted in black. [Figure](#page-5-0) 5a,b reveals several trends. First, increasing the Au concentration in NC Pt−Au systems leads to improved stability with respect to grain coarsening. For Pt-10 at. % Au systems, the crystalline grains depicted in [Figure](#page-5-0) 5a3,b3 are considerably smaller in size than those shown in [Figure](#page-5-0) 5a1,b1 for Pt-1 at. % Au. Second, in the small concentration limit shown in [Figure](#page-5-0) [5](#page-5-0)a1,b1 the Au concentration level is not sufficient to dope and subsequently pin all GBs ([Figure](#page-5-0) 5a1) or TJs [\(Figure](#page-5-0) 5b1). Third, a close examination of the non-dilute alloys with unrestricted segregation [\(Figure](#page-5-0) 5a2,a3) and the ones with TJ segregation only [\(Figure](#page-5-0) 5b2,b3) reveals that TJ segregation plays a dominant role in mitigating grain coarsening and that GB segregation provides a marginal improvement in the thermal stability of NC alloys.

The observations shown in [Figure](#page-5-0) 5a,b are quantified by tracking the temporal evolution of the average grain size ([Figure](#page-5-0) 5c) and total number of grains ([Figure](#page-5-0) 5d) in NC systems with Au concentrations of 1 and 10 at. % (see [Supplementary](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf) Figures S-3 and S-4 for grain coarsening results for [001] and [111] textures using all Au concentrations). Data for pure NC Pt are shown in [Figure](#page-5-0) 5c,d to provide a baseline for the comparison. Improved thermal stability is observed in both unrestricted and TJ segregation cases, where the increase in average grain size and the decrease in the total number of grains in these systems occur at rates slower than those in pure NC Pt. It is also interesting to note from [Figure](#page-5-0) 5c,d that TJ segregation is as effective in mitigating GB migration and grain coarsening in NC alloys as the cases with unrestricted segregation. For example, after 20 ns of annealing at 900 K, the difference in the average grain size between the system with TJ segregation and the one with unrestricted segregation is ∼0.5 nm for the Pt-10 at. % Au. The quantitative data in [Figures](#page-3-0) 3d and [5](#page-5-0)c clearly show that TJ segregation plays a major role in mitigating grain coarsening in NC alloys by locking the GB network and that GB segregation (i.e., unrestricted segregation cases) only provides a marginal improvement in the stability of these NC materials.

In summary, we performed large-scale atomistic simulations of solute segregation and high-temperature thermal annealing to examine the impact of TJ segregation on grain coarsening in a model NC Pt−Au alloy. To delineate contributions to the stability of nanostructures due to TJ segregation from those due to GB segregation, we explored two scenarios: unrestricted segregation where Au can segregate anywhere in the NC systems, and TJ segregation, where Au segregation is confined to TJ regions. While simulated systems of unrestricted segregation are more representative of experimentally observed segregation behavior in NC alloys, our computational studies of TJ segregation served to quantitatively describe the relative role of doped TJs in the stability of NC alloys. The NC systems were thermally annealed at 900 K, and the evolution of the grain structures was quantified by tracking the average grain size and number of grains as a function of time. It was shown that Au segregation to TJs renders these line defects immobile, thereby locking the GB network and hindering boundary migration. In dilute alloys, both unrestricted and TJ segregation cases are not as effective in mitigating grain coarsening, as the solute concentration level is not sufficient to dope and pin the GBs or TJs. Our work unequivocally shows that TJ segregation dominates the stability of NC alloys with respect to grain coarsening, and it highlights the need to

account for TJ effects in efforts aimed at discovering novel chemistries to fabricate thermally stable NC alloys. Finally, a natural extension of this work would be to explore the interplay among TJ segregation, initial grain size, annealing temperature, and the net impact on the thermal stability of NC alloys.

■ **ASSOCIATED CONTENT**

\bullet Supporting Information

The Supporting Information is available free of charge at [https://pubs.acs.org/doi/10.1021/acs.nanolett.4c02395.](https://pubs.acs.org/doi/10.1021/acs.nanolett.4c02395?goto=supporting-info)

Derivation of the triple junction to grain boundary density $\rho_{\text{TI}}/\rho_{\text{GB}}$, details on sample preparation and structural and chemical characterization of nanocrystalline Pt−Au alloys, and grain coarsening results for [001] and [111] nanocrystalline Pt−Au alloys using Au concentrations of 1, 5, and 10 at. % [\(PDF\)](https://pubs.acs.org/doi/suppl/10.1021/acs.nanolett.4c02395/suppl_file/nl4c02395_si_001.pdf)

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Notes

The authors declare no competing financial interest.

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